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(54) SULFOSUCCINATE DERIVATIVES FOR USE AS DETERGENT BUILDERS

(71) We, UNILEVER LIMITED, a company organised under the laws of Great Britain, of Unilever House, Blackfriars, London, E.C.4, England, do hereby declare the invention for which we pray that a patent may be granted to us and the method by which it is to be performed, to be particularly described in and by the following statement:—

Eutrophication is the process of excessive fertilization of aquatic plants through enrichment of waters with nutrients, such as carbon, nitrogen, phosphorus, potassium, iron, trace

metals and vitamins.

Although there is no present adequate proof, it has been postulated that the phosphoruscontaining builders present in detergent compositions can be a factor in eutrophication. Therefore any substitutes which do not contain phosphorus may decrease to some extent the eutrophication.

It is therefore an object of the present invention to provide novel compounds which are useful as detergent builders. It is another object of the present invention to provide 25 novel compounds which function as surface active agents and as detergent builders.

It has now been discovered that the alkali metal, ammonium and substituted ammonium salts of certain sulfoaliphatic dicarboxylic acids can serve as effective detergent builders in detergent compositions. Many of these detergent builders and their acid forms are novel and they are generally described as α - substituted - β - sulfosuccinic acids, and alkali metal, ammonium and substituted ammonium salts thereof, having the general formula:

wherein Z is O, S, SO, SO₂

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or NR_1 ; R is (C_1-C_{10}) alkyl, (C_1-C_{20}) hydroxyalkyl or (C_2-C_{20}) alkoxyalkyl; phenyl; carboxy-substituted or mono-, di- or tri-alkyl-substituted phenyl, wherein each of the alkyl group or groups contains 1-4 carbon atoms; sulfoalkyl or carboxyalkyl, wherein the alkyl moieties contain 1—4 carbon atoms; or R'Z(CH₂CH₂O)_n—CH₂CH₂—, wherein R' is hydrogen or alkyl containing 1—24 carbon atoms; and n is 0 or an integer of from 1-15; and when Z is

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or NR₁, the groups R may be joined with the nitrogen atom to form a morpholinyl group, and when Z is O and the group R is an alkyl group the latter contains 2-30 carbon atoms.

Thus, specific compounds and classes of compounds embraced by the generic formula

above include:

(I)

 α - alkoxy - β - sulfosuccinic acids α - phenoxy - β - sulfosuccinic acids α - carboxyphenoxy - β - sulfosuccinic acids

 α - alkylphenoxy - β - sulfosuccinic acids

 α - carboxyalkoxy - β - sulfosuccinic acids α - sulfosuccinic acids α - sulfosuccinic acids α - alkoxyethoxy - β - sulfosuccinic acids α - alkoxypolyethyleneoxyethoxy - β - sulfo-

succinic acids

 α - hydroxyalkoxy - β - sulfosuccinic acids;

the alkali metal, ammonium and substituted ammonium salts thereof; and the thio, sul-

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finyl and sulfonyl analogs of all the foregoing compounds wherein the oxygen group attached to the a-carbon of the succinic acid or succinate moiety is replaced by -S--SO- or -SO-, respectively, and/or wherein the cases of the a-alkoxyethoxy compounds and the a-alkoxypolyethyleneoxyethoxy compounds the oxygen attached to the alkyl group (R') is replaced by -S-, -SO- or _so,_.

In accordance with a specific aspect of the present invention are the nitrogen containing α - substituted - β - sulfosuccinic acids, and the alkali metal, ammonium and substituted ammonium salts thereof, having the following general formula:

$$R - N - CH - CH - SO_3H$$
 $R_1 - COOH COOH$
(II)

wherein the groups R and R1 may be the same or different and R1 is C1 to C20 alkyl, C₁ to C₄ hydroxyalkyl, carboxymethyl, carboxyethyl, sulfomethyl or sulfoethyl, or R and R₁ may be joined with the Nitrogen atom to form a morpholinyl moiety; and R1 is hydrogen or R.

Representative compounds and classes of compounds embraced by generic Formula II above include:

 α - alkylamino - β - sulfosuccinic acids such as α - methylamino, α - propylaimno, α - octylamino and α - laurylamino - β sulfosuccinic acid;

 α - dialkylamino - β - sulfosuccinic acids such as α - dimethylamino, α - ethylmethylamino, α - methylhexylamino and α - dioctylamino - β - sulfosuccinic acid;

 α - hydroxyalkylamino - β - sulfosuccinic acids such as α - hydroxyethylamino, α - hydroxybutylamino and α - bis(hydroxyethyl)amino - \(\beta \) sulfosuccinic acid;

 α - carboxyalkylamino - β - sulfosuccinic acids such as α - carboxymethylamino, α - carboxyethylamino - β - sulfosuccinic acid and the corresponding sulfo analogs;

 α - morpholinyl - β - sulfosuccinic

and the mono or poly salts thereof.

In accordance with further specific aspect of the present invention are the amine oxide derivatives of Formula II, wherein the nitrogen is a tertiary atom, corresponding to the general formula:

wherein the groups R may be the same or different and are as designated in Formula II.

As will be appreciated by those skilled in the art, the compounds of the invention contain at least two asymmetric carbon atoms and therefore can exist in several optically active forms as well as optically inactive mixtures (racemates). For purposes of this invention, the compounds as defined are intended to include all of the stereoisomeric forms and mixtures thereof.

In addition to the detergent building properties exhibited by the entire class of compounds described above, certain select members also exhibit properties which make them useful as wetting and foaming agents and thus constitute a class of novel surface active agents. For example, the α-alkoxy-β-sulfosuccinic acids and the thio analogs, containing from about 1-8 carbons, preferably from 1 to about 4 carbon atoms, exhibit excellent detergent building properties whereas the higher homologs containing from about 9-30 and more preferably 9-24 carbon atoms in the alkyl chain, additionally exhibit wetting, foaming and detergency properties.

Similarly, the α - alkowyethyl and aalkoxypolyethyleneoxyethoxy - β sulfosuccinic acid compounds containing from about 9-30 and preferably about 9-24 carbon atoms in the alkoxy moiety are also useful as wetting agents, foaming agents and detergents as well as detergent builders.

Although the builders of the present invention may be utilized as the free acid provided sufficient alkaline additives are included in the detergent composition to convert the acid forms in situ to the normal salt forms, the alkali metal, ammonium and substituted ammonium salts of the α-substituted-β-sulfosuccinic acids are preferred. Included in the substituted ammonium salts that can be employed are the monoethanolammonium, diethanolammonium, triethanolammonium, methylammonium, dimethylammonium, trimethylammonium, tetramethylammonium, morpholinium, N-methylmonoethanolammonium and N-ethylmonoethanolammonium salts and mixtures thereof.

The utility of the compounds of the present invention is not only reflected in terms of excellent building and biodegradability properties but also in low cost of preparation, since they are prepared from readily available and inexpensive materials. For example, the compounds of this invention are derived from sulfornaleic anhydride and readily available alcohols, thiols, hydroxy acids and amines.

More specifically, the compounds of this invention are reaction products derived by reaction between sulfomaleic acid and compounds having an active hydrogen atom.

Compounds having an active hydrogen atom and suitable for use in preparing the compounds of the present invention are monodi- or polyhydric alcohols and mono- di- or polyhydroxy acids and their sulfur-containing

analogs. Suitable examples of the aforementioned monohydric alcohols include alkoxyalkanols such as methoxyethanol and the linear primary and secondary alcohols containing up to 30 carbon atoms and their thio analogs; aromatic hydroxy compounds particularly the carbocyclic mono- and bicyclic aromatic hydroxy compounds, such as naphthols and phenols and the mono- di- or tri-C1-C4 alkyl ring substituted derivatives thereof. Suitable examples of dihydric alcohols include the glycols such as ethylene glycol, propylene glycol, butylene glycol, trimethylene glycol, tetramethylene glycol, pentamethylene glycol, hexamethylene glycol, heptamethylene glycol, long chain 1,2-diols containing from 8-30 carbon atoms and aromatic carbocyclic glycols such as phenylethylene glycol. Similarly, suitable polyhydric alcohols include glycerol, pentaerythritol, hexanetriol, sugars and their thio analogs.

In addition to the alcohols, the hydroxy carboxylic and sulfonic acids (in their ester and acid/salt forms, respectively) may also react with sulfomaleic anhydride and sulfomaleic acid. These include glycollic, lactic, glyceric, hydroxypropionic, salicyclic and mercapto acetic acid, hydroxymethanesulfonic acid and hydroxyethanesulfonic acid.

Still another important class of compounds containing active hydrogen are ethylene oxide adducts of C₁ to C₃₀ primary and secondary alcohols with 1—15 moles of ethylene oxide.

In general, the α - substituted - β - sulfosuccinate salts, wherein the α -substituent is joined to the α -carbon atom of the sulfosuccinate moiety by an O or S linkage, may be prepared by heating at a temperature of from about 25 to 120° C, preferably 60 to 100° C, sulfomaleic anhydride with a suitable compound having an active hydrogen atom as described above followed by further treatment with a base, for example an alkali metal hydroxide. The desired α -oxy or α -thio- β -sulfosuccinate may then be recovered and purified using conventional techniques.

The α - substituted - β - sulfosuccinate salts wherein the α -substituent is joined to the α -carbon atom of the sulfosuccinate moiety by an SO SO₂ linkage may be prepared by treating the appropriate α -substituted thio- β -sulfosuccinate with hydrogen peroxide according to the methods described on pages 471—472 in the text, "Reagents for Organic Synthesis" Fieser and Fieser, published by John Wiley & Sons, Inc., 1967.

The α - substituted - β - sulfosuccinate salts wherein the α -substituent is joined to the α -carbon atom of the sulfosuccinate moiety by an amino function (as in Formula II) may be prepared by reacting an appropriately substituted or unsubstituted primary or secondary amine with alkali metal salts of sulfomaleic acid. Typical amines suitable for reaction to

form the α - substituted amino - β - 6. sulfosuccinates include:

ethanolamine
diethanolamine
propanolamine
morpholine
N-methylethanolamine
glycine
alanine
N-methyl taurine
alkylamines containing 1—20 carbons in the
alkyl chain, as well as other amines having
a replaceable or active hydrogen and a
basicity comparable to the aforementioned
amines.

In particular the α-amino substituted-β-sulfosuccinates derived from water-soluble amines may be prepared by reacting in aqueous solution without the aid of heat and those derived from water-insoluble amines (i.e., higher alkylamines) are reacted in a mixed solvent system such as ethanol/water or dioxane/water at temperatures ranging from about 25° C to about 80° C; isolation from the reaction medium, and purification if desired, being effected by conventional methods.

The compounds wherein the α -substituent is joined to the α -carbon atom by an amine oxide group (as in Formula III) may be prepared by reacting the tertiary amine compounds of Formula II with oxidizing agents such as hydrogen peroxide, peroxyacetic and peroxyformic acid in the manner described for oxidizing tertiary amines by Hoh et al., J. Am. Oil Chemists' Soc., 40, 268 (1963).

In preparing the sulfosuccinate salts from the free acid, the amount of base utilized will determine whether the mono- di- or tri-salt is obtained. For example, the use of one mole of base (i.e., sodium hydroxide) per mole of α -hydroxy- β -sulfosuccinic acid yields the monosodium salt; the use of two moles of sodium hydroxide, the disodium salt and the use of three moles of sodium hydroxide, the trisodium salt. When R is carboxymethyl, carboxyethyl, sulfomethyl or sulfoethyl, a tetrasalt can also be obtained. Similarly, other bases, such as ammonium hydroxide and organic amines, may be utilized in the same manner to afford the type of salt desired.

According to the present invention, excellent cleaning results can be obtained by using the novel compounds described above as detergency builders with a wide range of detergent surface active materials and mixtures thereof in any of the usual physical forms for such compositions such as powders, beads, flakes, bars, tablets, noodles, liquids and the like. The builders can be used singularly, in combination with each other as the sole builder in the detergent composition or in combination

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with other well-known detergent builders such as sodium nitrilotriacetate, sodium ethylene-diaminetetraacetate, sodium tripolyphosphate, trisodium orthophosphate, sodium and potassium pyrophosphate, sodium polyacrylate, disodium oxydiacetate, trisodium citrate, trisodium carboxymethyloxysuccinate, salts of oxidized starches and sodium or potassium carbonate, as well as other conventional organic and inorganic builders.

When using the detergent compositions to wash clothes, the wash solutions should have a pH from about 7 to 12 and preferably from 9 to 11 throughout the washing cycle. Therefore, the presence of an alkaline buffer in the detergent composition is usually desirable particularly when the soil to be removed from the clothes has a high content of acidic components. Suitable buffers include any of the common organic and/or inorganic buffers such as monoethanolamine, diethanolamine, triethanolamine, sodium and potassium silicates, sodium and potassium carbonates and bicarbonates and the like.

In the detergent compositions, which are claimed in our copending divisional application No. 785/75 (Serial No. 1,398,422), the only essential ingredients are the detergent surface active material and the builder. The weight percent of the builder present in the detergent composition will range from 5 to 90% and preferably from 20 to 60% and more preferably 35—50% by weight of the total weight of the composition. When expressed as a weight ratio of builder to surfactant, the builders used in the instant invention will generally be present in a ratio of about 1:10 to about 10:1, and preferably 2:1 to 5:1 depending on the end use of whether a heavy-duty or light-duty detergent is desired. When the builders are used in mechanical dishwashing compositions, the ratio of builder to surfactant is from about 10:1 to about 50:1.

The detergent surface active compounds which can be used within the detergent compositions include anionic, nonionic, zwitterionic, ampholytic detergent compounds and mixtures thereof. A detailed description of such detergent compounds is to be found in the Specification of our Divisional Application referred to above.

Other materials which may be present in the detergent compositions in generally minor amounts are those conventionally present therein. Typical examples thereof include the well-known soil-suspending agents, hydrotropes, corrosion inhibitors, dyes, perfumes, fillers such as sodium sulfate, optical brighteners, perborates, bleaches, bleach activators, enzymes, suds boosters, suds depress-

ants, germicides, fungicides, anti-tarnishing agents, cationic detergents, fabric softening agents and in the case of liquid compositions, opacifiers and organic solvents. The balance of the detergent compositions may be water or inert filler.

It has been discovered that when higher than normal levels of anionic, nonionic, ampholytic or zwitterionic surfactants are used with the sulfosuccinate derivative salts of this invention, the detergency of the formulation is significantly enhanced particularly at low formulation concentrations (~ 0.1%) which are typically used by the housewife. For enhanced results the detergent formulation should contain surfactant levels of about 25% to about 45% by weight and the sulfosuccinate derivative salt levels of about 25% to about 75% by weight in the cases where the surfactants are anionic, ampholytic or zwitterionic. When the surfactant is a nonionic, enhanced detergency results are obtained when the level of said nonionic in the formulation is from about 15% to 30% by weight and the level of sulfosuccinate derivative salt is from about 25% to about 85% by weight.

In addition to their use in general household detergent compositions, the builders of the present invention find utility as boiler scale removers, stain removers and general chelating agents. When used at pH's of about 2 to about 5 as partially neutralized alkali metal, ammonium or substituted ammonium salts, especially in combination with wetting agents and surfactants, the compounds of the invention are excellent metal cleaning compounds.

EXAMPLE 1.

Table 1 below illustrates detergent compositions containing the detergency builders of the present invention in combination with representative classes of surface active agents compared with control or standard phosphatebuilt detergent compositions. The compositions were prepared by blending together the recited components in the proportions indicated, including an anticorrosive agent and buffer agent (sodium silicate), and the balance being water. The compositions were then tested on vacuum cleaner dust soiled cloth for detergency or cleaning ability in the Terg-O-Tometer test; wherein washing conditions are as indicated and the results reported as detergency units. The average detergency units (DU) of the formulation is the final reflectance of the soiled cloth, the reflectances being obtained by measurement with a Gardner automatic color difference meter, Model AC-3.

Component						ري ا	% Composition	Ton							ပိ	ntrols	Controls (% Composition)	sodu	tion)		
a-hydroxya	20	20	20	8	20															1	ļ
a-(2-hydroxy-ethoxy)a	æ					20	20														
a-dodecyloxya								20													
a-methoxya									20												
a-carboxymethoxya										. 02											
a-dodecyloxyethoxy ^a											20	•									
a-dodecylthioa												20									
Nas PaO10																			50	50	50
Sodium silicate (SiO ₂ :Na ₂ O=2.4:1)	10	10	10	01	10	01	10	01	10	, 9	10	10	10	10	10	01	10	10	10	10	10
Anionicb	18					18	36	18	18	18	18	18	82	81							
Anionicc		18													18				18		
Nonionicd			70													20				20	
Ampholytice				18													8		ı		00
Zwitterionicf					18				. 4	ميش								<u>~</u>) •
Water	1								```	- [_balance_	e e	İ								
Formulation Concentration, %	0.2	0.2	0.2	0.2	0.2 0.2 0.2 0.1 0.2	0.2	0.1	0.2	0.2	0.2	.0.2	0.2	0.1	0.2	0.2	0.2	0.2	0.2	0.2	0.2	0.2
Detergency (DU's)	24.6 24.4	24.4	26.5	23.1	26.5 23.1 25.6 26.1 24.7 27.1 23.0 24.2 15.4 22.3	26.1	24.7	27.1	23.0	24.2	15.4	22 3	•		7 7 7 7 7 7 7 7 7 7 7 7 7 7 7 7 7 7 7 7	9	•				

_6	1,39	98,421	6
5	 Sodium salts of α-substituted-β-sulfosuccinate Sodium linear secondary alkyl (C₁₀—C_{1s}) benzene sulfonate Sodium C_{1s}—C_{1e} α-olefin sulfonate C₁₁—C₁, linear secondary alcohols ethoxylated with 7 moles ethylene oxide/mole alcohol 	7.04 gm of sodium hydroxide in 50 ml water is then added and the mixture heated at 60° C for 2 hours. The heated mixture is then extracted three times with 300 cc portions of	65
10	methyl taurate) 'Sulfobetaine DCH (cocodimethylsulfopropyl betaine) Washing conditions—180 ppm (2:1 Ca++/Mg++); 120° F: pH 10 The following Examples 2 to 11 shows	fraction (at renux) and the acetone insoluble fraction is then filtered, washed with additional acetone and dried to give 16.2 gm of the title compound (structure confirmed by NMR and ion exchange of a sample followed by titration with standard sodium hydroxide: neutralization equivalent: found 147.0, theory 149.3).	70
15	the preparation of α -substituted- β -sulphosuccinates.	EXAMPLE 5.	75
20	EXAMPLE 2. Preparation of Trisodium α-thio substituted- β-Sulfosuccinate Sulfomaleic anhydride was prepared by heating a mixture of 1 mole of sulfur trioxide	Preparation of Trisodium c ₁ -Dodecyloxy- ethoxy-β-Sulfosuccinate Using the procedure of Example 4 and substituting 37 gm (0.32 mole) of 2-dodecyl- oxyethanol in place of the n-dodecanol there is obtained trisodium α-dodecyloxyethoxy-β- sulfosuccinate.	80
25	with 1 mole of maleic anhydride first at 60° C until the exothermic reaction subsided and then at 100—110° C for 3 hours. Forty-five gm of sulfomaleic anhydride product was then mixed with 45 gm of ice and the resulting solution extracted 25 times with 50 ml portions of either to remove maleic acid. A 20 ml	EXAMPLES 6—8. Using n-tetradecanol in place of n-dodecanol in the procedure of Example 4 and carrying out the hydrolysis step with 10% excess aqueous sodium hydroxide at 80° C for 4	85
30	collected, adjusted to pH = 8.6 with sodium hydroxide and refluxed for 4 hours. Then, an aqueous solution containing excess sodium methyl or ethyl mercaptide is added and the	hours instead of 60° C at 2 hours, there is obtained trisodium α-tetradecyloxy-β-sulfo-succinate (Example 6). Similarly, using n-hexadecanol in place of n-dodecanol, there is obtained trisodium α-hexadecyloxy-β-sulfo-succinate (Example 7); using n-octadecanol, there is obtained trisodium α-hexadecyloxy-β-sulfo-succinate (Example 7); using n-octadecanol,	90
35	resulting solution allowed to stand overnight. The reaction mixture is then passed through a column of cation exchange resin and the effluent to low volume. The pH is then adjusted to 8.5 with sodium hydroxide and the	sulfosuccinate (Example 8). EXAMPLE 9	95
40	solution evaporated to dryness. In this way, there is afforded trisodium α -methylthio- β -sulfosuccinate or trisodium α -ethylthio- β -sulfosuccinate.	Preparation of Trisodium c ₂ -Hydroxyethoxy- β-Sulfosuccinate Sulfomaleic anhydride (20 gm) is mixed with 37.2 gm (0.6 mole) of ethylene glycol and heated at 80° C for 4 hours. A solution of	100
45	EXAMPLE 3. Preparation of Trisodium α-Ethoxy-β- Sulfosuccinate Sulfomaleic anhydride, 3.9 gm, was dissolved in 25 ml of ethanol and refluxed for 5 hours. Then, 24 gm of 25% sodium ethoxide in ethanol was added and the	75 ml of water is then added and the mixture heated at 80° C for 4 hours. The mixture is then mixed with acetone to precipitate trisodium - α - hydroxyethoxy - β - sulfosuccinate, which is purified by repeated extractions with hot acetone to remove all ethylene glasses.	105
50	was then evaporated and the residue and	and water followed by filtration and drying in an oven.	110
55	dissolved in 100 ml of water and heated for 1 hour at 80° C. The solution was then decolorized with 5 gm of charcoal, filtered and evaporated. The crude residue of trisodium α -ethoxy- β -sulfosuccinate was purified by trituration with acetic acid and then filtered, washed with acetone and dried.	EXAMPLE 10. Preparation of the Tetrasodium Salt of α-Carboxymethyloxy-β-Sulfosuccinic Acid Sulfomaleic anhydride (19.4 gm) is heated to 60° C and combined with 76 gm of ethyl glycolate. After heating the mixture at 80° C for 5 1/2 hours, there is added a solution of	115
60	EXAMPLE 4. Preparation of Trisodium α-Dodecyloxy-β- Sulfosuccinate Sulfomaleic anhydride (10 gm) are mixed with 80 gm (0.45 mole) of n-dodecanol and	45 gm of sodium hydroxide dissolved in 135 gm of water and heating is continued at 80° for an additional three hours. The mixture is then diluted with water and passed through a column of a cation exchange resin. The resulting eluent is then evaporated in vacuo to	120

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a low volume and the concentrate extracted repeatedly with acetone to remove glycolic acid and other impurities. The resulting extracted residue is then dissolved in water and the pH of the solution adjusted to 8.6 with dilute sodium hydroxide. Evaporation of the solution in vacuo and drying over P2Os gives a residue of the tetrasodium salt of a-carboxymethyloxy- β -sulfosuccinic acid.

EXAMPLE 11.

Prepartion of Trisodium o-(2-Hydroxyethylamino)-β-Sulfosuccinate Sulfomaleic anhydride (prepared from a

1/1 ratio of SO₃/maleic anhydride by heating the mixture at 100—110° C for 3 hours; active content, 86%) 20.7 gm (0.1 mole), was added to 100 gm of ice. The resulting solution was kept at 10—15° C and neutralized to pH = 8.6 by slowly adding 28.4 gm of 50% sodium hydroxide. Monoethanolamine, 6.1 gm (0.1 mole), was next added slowly and the temperature allowed to rise to room temper-

ature. After standing overnight, the reaction mixture was poured into 800 ml of acetone. The solvent was then decanted from the resulting syrupy lower layer and the latter reprecipated three times from water with fresh acetone. The residue was then dried in a dessicator over P2O5 to give 38.8 gm of an off-white, granular product containing approximately 88% trisodium - α - (2 - hydroxyethylamino) - β - sulfosuccinate by tritration with perchloric acid. The structure was confirmed by NMR analysis.

The above method is satisfactory for reacting water-soluble amino compounds with sulfomaleic anhydride. For water insoluble amino compounds such as the higher alkyl amines, a mixed solvent system such as ethanol/water and dioxane/water is used together with temperatures in the range of 25-80° C.

Table 2 indicates the reactants required to obtain other α - substituted - β - sulfo-succinate salts having the R and Z moieties set forth in Formula I, following the procedures of the appropriate examples.

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		TABLE 2	
۳.	2	Exampl	Example Reactants (Sulfomaleic Anhydride +)
Ethyl	0	3	ethanol
Alkyl (C,-C,10)	0	4- 8-	C3-C30 primary, secondary or tertiary alcohol
Phenyl	0	4-8	Phenol
Carboxyphenyl	0	01	Methyl salicylate
(C,-C) Alkyiphenyl	0	4-8	(C ₁ -C ₄) Alkylphenol
o, m, p methylphenyl or methylisopropyl phenyls	0	4-8	o, m, p cresol, carvacrol, thymol
H 00C - CH,-	0	01	Ethyl glycolate
ноос-сн,сн,-	0	01	Methy I-B-hydroxypropionate
HO,S-CH,CH,-	0	4	Isethionic acid
R'O(CH,CH,O)n-CH,CH,-	0	4	R'O(CH,CH,O),CH,CH,OH
HOCH,CH,OCH,CH,-	0	4	носн,сн,осн,сн,он
но(с'н,сн,о),сн,сн,-	0	4.	но(сн,сн,о),сн,сн,он
HO(C'H,CH,O),OH,CH,-	0	ĭ.4	HO(CH,CH,O), CH,CH,OH
но(с'и,с'и,с'и,с'и,-	0	4	H0(CH,CH,0), CH,CH,0H
CH,CH,O(CH,CH,O),CH,CH,-	0	4	СН,СН,О(СН,СН,О),СН,СН,ОН
CH,(CH,),0(CH,CH,0),CH,CH,-	0	4.	CH,(CH,),,O(CH,CH,O),CH,CH,OH
CH,(CH,),,0(CH,CH,0),CH,CH,-	0	4	СН,(СН,),,0(СН,СН,О),СН,СН,ОН
CH,CH,S(CH,CH,CH,CH,-	0	य ∵	CH,CH,S(CH,CH,O),CH,CH,OH
CH, CH, S(CH, CH, O), CH, CH,	0	. 4	CH,CH,S(CH,CH,O),CH,CH,OH

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	TAB	TABLE 2 (Continued)	inued)
R	2	Example	Example Reactants (Sulfomaleic Anhydride +)
HOCH,CH,-	0	6	Ethylene glycol
носн,сңсн,-	0	6	Trimethylene glycol
HOCH,(CH,),CH,-	0	6	Hexamethylene glycol
Methyl or ethyl	S	7	Sodium methyl or ethyl mercaptide
Alkyl (C,-C,0)	S	. 4-8	C,-C,0) alkylmercaptan
Phenyl	S	48	Thiophenol
Carboxyphenyl	S	10	Methyl 2-mercaptobenzoate
Aikyiphenyi	S	4-8	Alkylthiophenol
H00C-CH ₂ -	S	10	Ethyl mercaptoacetate
ноос-сн,-сн,-	S	10	Methyl B-mercaptopropionate
HO,SCH,CH,-	S	, .4	Mercaptoethanesulfonic acid
R'O(CH,CH,O)nCH,CH,-*	S	4	R'O(CH2CH2O)nCH2CH2SH*
HOCH,CH,OCH,CH,-	S	2	HOCH, CH, OCH, CH, SNa
HO(CH,CH,O),CH,CH,-	છ	7	HO(CH,CHO),CH,CH,SNa
HO(CH,CH,O),0CH,CH,-	S	2	HO(CH,CH,O),OH,CH,SNa
HO(CH ₂ CH ₂ O) ₁₅ CH ₂ CH ₂ —	S	2	HO(CH,CH,O), GH,CH,SNa
CH1CH1O(CH1CH1O)1CH1CH1-	S	4	CH,CH,O(CH,CH,O),CH,CH,SH
CH,(CH,),1,0(CH,CH,O),CH,CH,-	S	. 4	CH,(CH,),,(CH,CH,O),CH,CH,SH
CH,(CH,),,0(CH,CH,0),CH,CH,-	Ø	4	CH,(CH,),,0(CH,CH,O),CH,CH,SH

	TABLE	TABLE 2 (Continued)	ued)
~	7	Example	Example Reactants (Sulfomalcic Anhydride +)
CHCHS(CH,CH,O),CH,CH,-	S	4	CH,CH,CH,CH,SH
CH,CH,S(CH,CH,O),CH,CH,-	S	4	CH,CH,S(CH,CH,O),CH,CH,SH
носн,сн,-	S	7	Sodium 2-hydroxyethylmercaptide
носн,си,сн,–	S	· ~	Sodium 3-hydroxypropylmercaptide
HOCH,(CH,),CH,-	S	7	Sodium 6-hydroxyhexylmercaptide
Methyl or ethyl	80	2	Methyl or ethyl mercaptan ^a
Alkyi (C,-C,0)	80	4 8-8	Cs-C30 alkylmercaptana
Phenyl	SO	4-8	Thiophenol ^a
Carboxyphenyl	SO	10	Methyl 2-mercaptobenzoatea
C ₁ C ₄ alkylphenyl	SO	4-8	Alkylthiophenol ^a
ноос-сн,-	SO	10	Ethylmercaptoacetatea
ноос-сн,сн,-	SO	10	Methyl <i>B</i> -mercaptopropionate ^a
R' O(CH2CH2O)nCH2-*	SO	S	R'O(CH,CH,O),CH,CH,SHa *
HOCH, CH, -	80	7	Sodium 2-hydroxyethylmercaptide
носн,сн,сн,~	SO	7	Sodium 3-hydroxypropylmercaptide
HO(CH,CH,),CH,-	80	7	Sodium 6-hydroxyhexylmercaptide
a Followed by oxidation to sulfoxide according to Fieser reference.	ser refere		
C,-C, alkyi	so,		C,-C, alkylmercaptanb
C,-C,0 alkyl	so,	4-8	C ₃ C ₃₀ alkylmercaptan ^b

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R	2	Example	Example Reactants (Sulfomaleic Anhydride +)	
Phenyl	SO2	4-8	A-8 Thiophenolb	
Carboxyphenyl	SO,	4-8	4-8 Methyl 2-mercaptobenzoateb	
H00C-CH ₂ -	SO ₂	10	Ethyl mercaptoacetateb	
H00C-CH,CH,CH,-	SO ₂	10	Methyl 2-mercaptobutyrateb	
R' 0(CH,CH,CH,CH,CH,-*	502	. 2	R'O(CH,CH,O)nCH,CH,SHb*	
b Followed by oxidation to sultone according 'o Fieser reference.	er refere	ıce.		

* wherein R, is hydrogen or a C,-C, alkyl group.

wherein R' is hydrogen or alkyl containing 1-24 carbon atoms and n is 0 or an integer of 1-15; and Z is O, S, SO₂, SO, WHAT WE CLAIM IS:—
1. An α - substituted - β - sulfosuccinic acid having the general formula:

 $R-Z-CH-CH-SO_3H$

соон соон

S

or NR, wherein R, is hydrogen or R as defined above; when Z is

or an alkali metal, ammonium or substituted

2

or NR, the groups R may be joined with the nitrogen atom to form a morpholinyl group, and when Z is O and the group R is an alkyl group, the latter contains 2—30 carbon atoms.

ammonium salt thereof; wherein R is (C.—C.a.) alkyl, (C.—C.a.) hydroxyalkyl or (C.—C.a.) alkoxyalkyl; phenyl; carboxysubstituted or mono-, di- or tri-alkyl-substituted phenyl wherein each of the alkyl groups contains 1—4 carbon atoms; sulfoalkyl or carboxyalkyl wherein the alkyl moieties con-

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R'Z(CH,CH,O),—CH,CH,—

15

tain 1-4 carbon atoms; or

A compound as claimed in claim 1 wherein Z is oxygen.
 A compound as claimed in claim 1 or claim 2 wherein R is an alkyl group containing
 2-24 carbon atoms.
 A compound as claimed in claim 2

4. A compound as claimed in claim 3 wherein R is an alkyl group containing 9—24 carbon atoms.

 A compound as claimed in claim 3 wherein R is an alkyl group containing 2—8 carbon atoms.

6. α - dodecyloxy - β - sulfosuccinic acid or an alkali metal, ammonium or substituted ammonium salt thereof.

7. α - hexadecyloxy - β - sulfosuccinic acid or an alkali metal, ammonium or substituted ammonium salt thereof.

8. α - octadecyloxy - β - sulfosuccinic acid or an alkali metal, ammonium or substituted ammonium salt thereof.

9. An α -hydroxyalkoxy (2—24 carbon atoms)- β -sulfosuccinic acid or an alkali metal, ammonium or substituted ammonium salt thereof.

25 10. A hydroxyethoxy compound as claimed in claim 9.

 An α-alkoxy (1—24 carbon atoms)ethoxy-β-sulfosuccinate as claimed in claim 1.

α - dodecyloxyethoxy - β - sulfosuccinic acid or an alkali metal, ammonium or substituted ammonium salt thereof.

13. A compound as claimed in claim 1 wherein R is a carboxymethyl group.

14. A compound as claimed in claim 1 35 wherein Z is sulfur.

15. A compound as claimed in claim 14 wherein R is an alkyl group containing 1—24 carbon atoms.

16. A compound as claimed in claim 14

wherein R is an alkyl group containing 1—8 40 carbon atoms.

17. α - dodecylthio - β - sulfosuccinic acid or an alkali metal, ammonium or substituted ammonium salt thereof.

18. α - hexadecylthio - β - sulfosuccinic acid or an alkali metal, ammonium or substituted ammonium salt thereof.

19. α - octadecylthio - β - sulfosuccinic acid or an alkali metal, ammonium or substituted ammonium salt thereof.

20. A compound as claimed in claim 1 wherein Z is SO.

21. A compound as claimed in claim 1 wherein Z is SO₂.

22. A compound as claimed in claim 1 55 having the general formula:

$$R - N - CH - CH - SO_3H$$
 $R_1 - COOH COOH$
(II)

wherein the groups R and R₁ may be the same or different, and R is C_1 — C_{20} alkyl; C_1 — C_4 hydroxyalkyl; carboxymethyl; carboxyethyl; sulfomethyl; sulfoethyl; or R and R₁ may be joined with the nitrogen atom to form a morpholinyl moiety; and R₁ is hydrogen or R.

23. A compound as claimed in claim 1 substantially as described herein with reference to any of Examples 2—11 and in Table 2.

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